AMENDMENT UNDER 37 C.F.R. § 1.111 Attorney Docket No.: Q80857

Application No.: 10/592,007

REMARKS

Claims 1 to 9 are all the claims pending in the application, prior to the present

Amendment.

Applicant has amended claims 8 and 9 to place them in independent form. Claim 9

contains recitations from claims 1 and 2. Claim 8 contains recitations from claims 1, 2 and 4.

Applicant has also added a new independent claim 10 that contains recitations from claims 1, 2

and 5. In addition, applicant has added new claims 11 and 12 that depend from claim 8 and

contain recitations from claims 6 and 7. Claims 1-7 have been canceled.

Applicant has amended the present specification to place it in better form and to correct a

number of obvious errors.

Claim 6 has been objected to because the formula contains a "y" subscript, but the

specification only describes an "x" subscript.

Applicant has canceled claim 6. Accordingly, this rejection is moot.

In general, applicant points out that the present specification contains a reference and

description of "y" at numerous locations, such as at pages 7, 9, 11, 12, 13 and 14. For example,

the present specification at page 13, lines 33-34 discloses that, in general, x = y, but "x" may be

different from "v." New claim 11 contains a "y" recitation.

Claim 2 has been rejected under the second paragraph at 35 U.S.C. §112 as indefinite.

The Examiner states that the term "parts" is a relative term which renders the claim

indefinite.

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Applicant has canceled claim 2. Accordingly, this rejection is moot with respect to claim 2.

Applicant notes that the term "parts" is employed in amended claims 8 and 9, and new claim 10. These claims state that the palladium element is present in an amount of from 0.001 to 15 parts by mass based on 100 parts by mass of $W_aZ_rO_{x_t}$ as set forth in the present specification at page 6, lines 33-35 and page 8, lines 13-16.

Claims 1, 2, 6 and 7 have been rejected under 35 U.S.C. §102(b) as anticipated by or, in the alternative, under 35 U.S.C. §103(a) as obvious over EP 1 063 010 to Ohtsuka et al.

Applicant has canceled claims 1, 2, 6 and 7.

Accordingly, applicant submits that this rejection is moot.

Claims 1, 2, 6 and 7 have been rejected under 35 U.S.C. §102(b) as anticipated by or, in the alternative, under 35 U.S.C. §103(a) as obvious over U.S. Patent 5,902,767 to Kresge et al.

Applicant has canceled claims 1, 2, 6 and 7.

Accordingly, applicant submits that this rejection is moot.

Claims 1, 2-5, 8 and 9 have been rejected under 35 U.S.C. §102(b) as anticipated by or, in the alternative, under 35 U.S.C. §103(a) as obvious over EP 0 620 205 to Suzuki et al.

Applicant has canceled claims 1 and 2-5, thus leaving amended claims 8 and 9, and new claims 10, 11 and 12 as being subject to this rejection.

Applicant submits that EP '205 does not disclose or render obvious the subject matter of claims 8-12 and, accordingly, requests withdrawal of this rejection. Attorney Docket No.: Q80857

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Claims 8-10 as set forth above are independent claims that are directed to a process for producing an oxygen-containing compound by reacting an olefin and oxygen in a gas phase in the presence of a catalyst represented by the formula Pd/W_aZrO_x. In this formula, Pd is a palladium-containing compound, a is a W/Zr molar ratio of from 0.01 to 5.0, and x is a value defined by the oxidized state of tungsten (W), zirconium (Zr) and palladium (Pd), and wherein the content of palladium element in the catalyst is from 0.001 to 15 parts by mass based on 100 parts by mass of WaZrOx.

Claim 9 is directed to producing acetic acid by reacting ethylene and oxygen in the presence of the catalyst. Claim 8 is directed to reacting propylene and oxygen and claim 10 is directed to reacting a butene and oxygen, to produce at least one compound selected from a number of different oxygen-containing compounds.

EP '205 discloses a catalyst for producing acetic acid by reacting ethylene and oxygen. The catalyst comprises metallic palladium and one or more hetero poly-acids that may contain one hetero-atom and one or more poly-atoms. EP '205 does not disclose any example of a catalyst containing the components Pd/Zr and W. Moreover, EP '205 does not disclose or suggest a process that employs the catalyst set forth in claims 8-10.

It is known that there are two types of heteropolyacids: Keggin-type and Dawson-type. See Fig. 1 on page 2 of the enclosed paper: Makoto Misono, "Unique acid catalysis of heteropoly compounds (heteropolyoxometalates) in the solid state," Chem. Commun., 2001, 1141-1152. When a heteropolyacid contains Zr and W, the composition of the heteropolyacid is ZrW₁₂O₄₀⁴or Zr₂W₁₈O₆₂8-. In other words, the W/Zr molar ratio of heteropolyacids is 12 or 9, which falls outside of the scope of the catalyst set forth in claims 8-10, wherein the W/Zr molar ratio is from

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0.01 to 5. Thus, EP '205 does not disclose or suggest a process that employs the catalyst set forth in the present claims.

In addition, EP '205 to Suzuki et al is the EP counterpart of Patent Document 6

(JP 7-89896 A) described in Background Art of the Description of the present application. In

Comparative Example 3 of the present application, as disclosed at page 19 of the present
specification, Catalyst 11 was produced according to JP 7-89896. In Comparative Example 6 of
the present application, an ethylene oxidation reaction was performed by using Catalyst 11. As
shown in Table 3 on page 21 of the present application, the acetic acid selectivity and the space
time yield (STY) achieved by employing the catalyst set forth in the present claims are greater
than that obtained by using the Catalyst 11 produced according to JP 7-89896, which
corresponds to EP '205 to Suzuki et al. Thus, EP '205 does not obtain the results achieved by
the present invention.

Further, EP '205 does not disclose or suggest a process employing propylene as set forth in claim 8 or a butene as set forth in claim 10.

In view of the above, applicant submits that EP '205 to Suzuki et al do not disclose or suggest the process set forth in the present claims or the results obtained by the present invention and, accordingly, requests withdrawal of this rejection.

In view of the above, reconsideration and allowance of this application are now believed to be in order, and such actions are hereby solicited. If any points remain in issue which the Examiner feels may be best resolved through a personal or telephone interview, the Examiner is kindly requested to contact the undersigned at the telephone number listed below.

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The USPTO is directed and authorized to charge all required fees, except for the Issue Fee and the Publication Fee, to Deposit Account No. 19-4880. Please also credit any overpayments to said Deposit Account.

Respectfully submitted,

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Unique acid catalysis of heteropoly compounds (heteropolyoxometalates) in the solid state

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Fundamental and superior characteristics of heteropoly compounds (heteropolyoxometalates) in the solid state that make them suitable for catalyst design at the atomic/molecular levels are described, together with important principles required for the understanding and design of solid heteropoly catalysts. First, the molecular nature of heteropolyanions (metal oxide clusters), which can be preserved in the solid state, enables control of the acid and redox properties over a wide range. Second, the presence of hierarchical structures (primary, secondary and tertiary structures) can lead to three catalysis modes—surface-type, pseudoliquid (or bulk-type I) and bulk-type II. Precise control of pore size is possible through the understanding of the microstructure, which results in unique shape selectivity observed for various reactions. Heteropoly compounds are green catalysts functioning in a variety of reaction fields and efficient bifunctional catalysts when comblined with other components. The elucidation of catalytic processes is also possible at the atomic/molecular level due to their molecular nature. The positions and dynamic nature of protons as well as organic reaction intermediates in the pseudoliquid phase can be clarified by spectroscopic techniques. Various reactions promoted by solid heteropoly catalysts are collected from recent publications to illustrate the usefulness of the above ideas and of heteropoly catalysts themselves.

1 Catalyst design at the atomic/molecular level 1.1 Importance and feasibility

Efficient catalysts are key materials in chemical technologies which supply useful substances to society and assist maintain-

Makoto Misono received his Bachelor, Master, and Doctor's degree in Engineering from the University of Tokyo and started his avademic career in 1966 at the University of Tokyo, where he hecame a full professor in 1983. He spent two years in the USA for his postdoctorate (1967-1969: at the University of California, Santa Barbara, and Mellan Institute, Pittshurgh). He has been studying heterogeneous catalysis mainly of mixed usides for 40 years and gave a plenary lecture at the 10th international Congress on Catalysis, Budapest, in 1992 on the subject of molecular catalyst design of solid heteropoly compounds. Since his retirement from the University of Tokyo with the Emeritus professorship in 1999, he has been a professor at Kogakuin University, a private technical university located in the center of Tokyo. He is a former president of Cetalysis Society and a vice president of Chemical Society of Japan. He was elected a member of Science Council of Japan in 2000. He is now more involved in various activities related to the environment and chemistry such as green/sustainable chemistry and clean fuel-exhaust of automobiles.

ing the environment as healthy as possible; in short, catalysis represent a key technology for a sustainable acciety. Although recent progress has been remarkable for homogeneous and blochemical canlysts, in particular the latter, heterogeneous consideration and the state of th

It is much more difficult to design heterogenous caralysis than to interpret reactions over mode solid catalysis. Hence, the development of practical (or commercial) catalysis still mostly relief on trial-and-error approaches, assisted by phenomenological knowledge on existing commercial catalysis and sophisticated knowledge on simple model catalysis. The design of practical catalysis and the atomic or molecular level has long been pursued. 1-3 since precisely designed multifunctional catalysis are obviously designed multifunctional catalysis are obviously designed multifunctional to this level is still a distant goal in many instances.

1.2 Our approach

We have endeavored in the last two decades to establish the methodology of catalyst design by studying crystalline mixed oxides. In addition to their crystallinity, our criteria to choose catalyst materials are; (i) wide variation of composition whilst retaining the basic crystal structure and (ii) catalytic performance near the level required for practical application. Such materials include heteropoly compounds, perovskite-type mixed oxides and zeolites whose structures can be well defined at least for the solid bulk phases. Differences between the surface and solid bulk phase appear to be much smaller for these metal oxides than for metallic catalysts. This is particularly true for heteropoly compounds if the preparation and characterization of catalysts are carefully carried out. Here, the term 'heteropoly compounds' (abbreviated as HPAs) will be used for heteropolyoxometalates which include heteropolyacids and their derivatives. Heteropolyacids are hydrogen forms of heteropolyanians produced by the condensation of more than two kinds of executions. Typical heteropolyanions are shown in Fig. 1. Using heteropoly compounds we attempted to establish the following relationships at the atomic/molecular levels as shown in Scheme 1.



This article reports important progress to establish the basis for catalysts design since the publication of special issue of Chemical Reviews on polyocometalates in 1998.3 It will lattempt to show that heteropoly compounds have various attractive and important characteristics in terms of catalysis and are promising materials for catalyst design at the atomic/molecular level.

DOI: 10.1039/6102573m

Chem. Commun., 2001, 1141, 1152, 1141

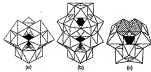


Fig. 1 Examples of heteropolyanions; (a) Keggin-type polyanion e.g. α -PW₁₂O₄₀¹·· (b) Dawson-type polyanion e.g. $P_2W_{18}O_{62}^{a}$ ·· (c) disubstituted polyanion e.g. γ -Si W_{10} Fe₂O₄₀¹⁰··.

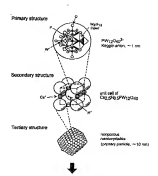
Cutilysts based on heteropoly compounds are hereafter denoted heteropoly (or HPA) catalysts. Earlier books and review articles may be referred to regarding the basic chemistry* and general catalysis* of HPAs. HPA catalysts have already been applied to several large-scale commercial processes.*

2 Basic concepts unique for solid HPA catalysts

In my view, the following concepts are essential to understand and design HPA catalysts, in addition to the knowledge generally required to understand heterogeneous catalysis of the ordinary mixed oxides.

2.1 Primary, secondary and fertiary structures

At an early stage of our study it was recognized that the hierarchical structure of solid HPAs was important for the understanding of the beterogeneous HPA catalysis, and we denoted the substructures as primary, secondary and tertiary.7 This may appear a very simple idea, but enormously helped the progress of our research, Fig. 2 shows a simplified illustration of such hierarchical structures. The primary structure is the structure of heteropolyanion itself i.e. the metal oxide cluster molecule and details the molecular nature of solid HPA catalyst Fig. 2 shows the most ubiquitous form which has the Keggin structure. The secondary structure is the three-dimensional (usually regular) arrangement consisting of polyanions, counter cations and additional molecules. The secondary structure is flexible to different extents depending on the counter cation and the structure of the polyanion, and is the basis of bulk-type catalysis of solid HPA catalysts (see below). The tertiary structure represents the manner in which the secondary structure assembles into solid particles and relates to properties such as particle size, surface area, and pore structure, and plays an important role in heterogeneous catalysis. Without understanding this hierarchical structure, one can neither understand solid HPA catalysts properly nor take advantage of their molecular nature, and the idea of bulk-type catalysis described below would not have evolved.



Porous eggregate of nanocrystatites (secondary particle, 0.1 -- 0.5 µm)

Fig. 2 Primary, secondary and tertiary structures; hierarchical structure of heteropoly compounds (HPAs) in the solid state.

2.2 Three types of catalysis

We have demonstrated that there are three totally different modes of catalysis for solid HPAs (Fig. 3), Surface-type catalysis (a) is ordinary heterogeneous catalysis which takes place on the solid surface (two-dimensional reaction field on outer surface and pore wall). Modes (b) and (c) represent bulktype catalysis where the reaction fields are three-dimensional in contrast to the surface-type catalysis. When the diffusion of resutant molecules in the solid (diffusion in the lattice rather than pores) is faster than the reaction, the solid bulk forms a pseudo-liquid phase in which catalytic reaction can proceed Fig. 3(b)]. In this instance, reactant molecules in the gas or liquid phase penetrate in between the polyanions (primary structure), sometimes expanding the distance between the polyanions, and react inside the bulk solid. The products come out to the surface and are released to the gas or liquid phase. Pseudoliquid catalysis, proposed in 1979, was not favourably accepted initially, since it appeared very unusual for heterogeneous mixed oxide catalysts. Now, however, such catalysis is more firmly established. So-c In the pseudoliquid phase such catalysts appear as solids but behave like liquids (solvent). As the active sites in the solid bulk e.g. protons, take part in catalysis, very high catalytic activities are often observed in the bulk phase. Phase transitions accompanied by an abrupt change in catalytic performance are also observed.50

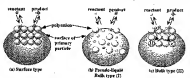


Fig. 3 Three types of catalysis for solid heteropoly compounds; (a) surface type; (b) pseudoliquid; butk type (I), (c) bulk type (II).

The second bulk-type catalysis (bulk-type (II), Fig. S(c)) is found for oxidation catalysis at high temperature when the diffusion of redox carriers (protons and electrons in this case) is rapid in the solid bulk, and the whole bulk participates in the reduction-oxidation cycle? It should be noted that, as the contribution towards catalysis of the solid bulk, varies with the relative rate of diffusion to that of reaction, intermediate cases between surface and bulk canalysis arise. Solid IFTA somaning cations of low ionic radii to charge ratio (If*, Ner., Cab*, etc.) between surface and bulk canalysis arise. Solid IFTA containing cations of low ionic radii to charge ratio (If*, Ner., Cab*, etc.) proceedings the service of the containing of the containing of the salts (searcely soluble in water, due to low solvation energy) usually show only surface-type catalysis.

The importance of this idea may be clearly understood by the following examples. Fig. 4(a) plots the relative activity and number of strong acid sites us, the extent of neutralization (or the Na content) for acidic Na salts of H₂PW₁₂O₄₀, ¹⁰ Hereafter, H₃PW₁₂O₄₀, one of the most widely used HPA catalysts is abbreviated as HPW unless stated otherwise. The reactions are of bulk type (I) and the acidity is related to the number of protons in the entire bulk. It is evident in Fig. 4(a) that the rates of bulk-type reactions and the bulk acidity decrease monotonically with the Na content, thus showing good parallelism between catalytic activity and acidity. This also demonstrates that the acid catalysts can be designed by control of their acidity in this manner. In contrast, no monotonical change is found for several reactions catalyzed by acidic Cs salts, Cs.H1 .PW :> O40 (denoted CsX), as shown in Fig. 4(b).54.6.11 These reactions are of surface type, as revealed by plots of the rates vs. she surface acidity (number of protons on the surface). The surface area sharply rises from 1-2 m2 g-1 for Cs1 and Cs2 to ca. 150 m2 g-1 for Cs3. The mechanism of the increase in surface area is interpreted in a later section of the article. As is obvious in Fig. 4(c), a good correlation is obtained.12 Another important point to be noted is that the specific activity of CsX [entalytic activity per surface proton, the slope in Fig. 4(c)] is much higher than known solid suids such as zeolites and silica-alumina. This fact demonstrates the high performance of HPA catalysts.

Bulk-type catalysis (1f) is also an essential concept required to understant and design HPA catalysts. ¹This bas been found to be relevant for oxidation reactions at high temperatures, I find the relevant for oxidation reactions at high temperatures, I find the set of bulk-type oxidation (explore) as a plotted against the surface redox property, very poor correlations are found, but the rates exhibit good correlations with bulk redox property. In contrast, the rate of surface-operation correlations very well with the surface redox property. Oxidation correlates very well with the surface redox property. Without knowing this fact, the development of practical coldation correlations would be surface.

2.3 Merits of HPA catalysts

The advantages of hetropoly compounds for heterogeneous catalysts are summarized in Table 1.5 There are several large-scale industrial processes utilizing HPA catalysts as oxidation and acid catalysts both in the solid state and in solution. **AAA* Most are environmentally friendly, so that HPA catalysts are reparted as promising green (or sustainable) catalysts. **ISA**

Table 1 Merits of solid heteropoly establish for catalyst design at the atomic/molecular level

- Systematic variation of said and redox properties are possible for entailors design.
- Molecular nature of solid heteropoly employeds originating from heteropolyanion molecules enables precise design of catalysts and molecular description of catalysts processes
- A variety of reaction fields are available for establic systems.

 The unique basicity of polyanions

3 Novel aspects of pseudoliquid catalysis

-3.1 Variety of reaction fields

Pseudoliquid behavior, Dulk-tyne (f) estalysis) has also bost found for liquid-soil beteregeneous systems. For example, the relative catalytic activities of HPW and Cs2.3 dramatically change with the polarity of resource generation (Cs2.3 for pinacol rearrangement (pseudoliquid catalysis) and HPW < Cs2.5 for allysistion of aromatica (surface-type catalysis). HPW shows high activities for the reactions in the solid bulk (pseudoliquid), since protons in the behavior of the control o

The earlysis of solid HPAs in the figuid phase has been well documented. He or example, three sections difficulty in polarity of reactasts were compared using several skall- and sklating- earth salts of HPW. It was shown that the decomposition of exclusive street is catalyzed on the solid surface fourface, the proposition of prison of the surface fourface is catalyzed on the solid surface fourface the place of the proposition of the president process of the present skill of the present skill of the present of the present street is made to the present solid state. The solid state is made button-to it mainty catalyzed by HPA disadved in its solid state. The order of catalytic activity, therefore, is very different between the reactions. This again demonstrates the importance of distinguishing surfaces and bull-three catalytics for the catalysts of the catalysts.

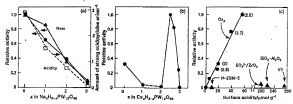


Fig. 4 Importance of differentiating meface- and bulle-type analysis, (a) Bull-type (1) pseudobjund) catalysis both both, acidiny of No store $I(I)/W_1/G_{10}$ (IMPN): (4) Conversion of melabous, (4) elwhystime of propose-2c. and Cli bell-worldy measured by exemply hold profiles, (b) Rates of silvagues of 1.3-5 trimothylentone by cyclohexner catalyses by Crashin of HPW (meta-cype catalysis), (c) Rates of alsylation over Cc salts (data from Fig. 4(n)) are plotted in the meta-catalysis operation of the catalysis operation of the conversion three colors.

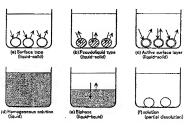


Fig. 5 Various reaction fields of HPA catalysis in reaction systems containing a liquid phase.

design of HPAs, Fig. 5(a) and (d) correspond to the ordinary heterogeneous camitys fit(juil could) and homogeneous (solution) cetalysis, respectively. Phase-mansfer caulysis for liquid-injuid biphase systems [Fig. 5(c)] is well known and there are at least two large-scale commercial processes using HPA catalysts. Are Catalysis by solid HPAs for solid states reactions 1 and in a new phase formed on the surface layers [Fig. 5(c)] are new. In the latter, protons introduced on the surface layer of C33 create a new active phase similar to a "pseudoliquid" (Fig. 6). Among other recent interesting observations is the using well as the using the control of the contr

relationship between the shape of the primary structure, the resulting secondary structures, and the remarkable influence on the estalytic activity.18 The thermodynamically favored synthesis of methyl terr-butyl ether (MTBE) from isobutylene and methanol proceeds at low temperature (323 K) on Dawson-type hetropolyscids (HeP2W1AOe2 and HeP2Mo18Oe2) repidly and selectively. In contrast, the reaction is very slow over Keggintype heteropolyacids, in spite of the higher acid strength and comparable acid content. We found that Dawson-type heteropolyacids are amorphous under the reaction conditions due to the ellipsidal shape of the polyanion, whereas Keggin-type polyanions having spherical shape are crystalline (bec structure). Owing to this difference the former forms active pseudoliquids, while the latter are much less active (Table 2). As for MTBE synthesis catalyzed by H4SiW12O40, Blelanski's group has made an extensive study and proposed a hypothesis that the reaction takes place on the surface to which methanol and protons are supplied from the pseudoliopid phase, 19

3.2 Protons, water and organic molecules in pseudoliquids

To understand and destijan pseudoligaid catalysts at the molecular level, information about the actific promos, such as their location, mobility and donating ability (acid strength), is indispensable. Information about the insusations between actific protons and small basic molecules such as water and actions provides such this knowledge about pseudoligaid catalysis, since water is often contained in the working state and plays an important to lee. Furthermore, the states and whamies of protons in the solid arm interesting subjects of solid-state chemistry. In our early studies we observed protons in HPW using MAS NMR, then directly detected by a combination of NMR and RI (it he reaction intermediates of ethanol dehydration in the pseudoliquid phase, and disclosed the dynamic behavior of metanol.**

While IR is usually a powerful tool to study solid catalysts the IR spectra of OH bands of heteropolyacids are ambiguous, For highly hydrated HPW, the OH stretching and bending modes of





Fig. 6 Catalysis in a rowel phase formed on the surface layer of a solid HPA for a liquid-solid reaction system. Hydrothysis of bittineshiple propue monodistantal in swater at 248 K. The receiver rate rises sharply when the second component is added after a certain reaction time (incleaned by arms). The reace she're the addition are much higher than the user of the rates of each continuous, indicating the formation of an active layer on the oriface of CS3.

Table 2 Unique relationships between shope of polyanion (primary structure), secondary structure and estalytic activity

Primary structure* (shape)	Secondary structure at the working state	Catalytic activity for MTBE synthesis at 323 K	
Dawson (allipse) HaP2W14O42 HaP2MU14O42	Amorphous	Very high	
Keggin (spherical) H ₂ PW ₁₂ O ₄₀ H ₄ SiW ₁₂ O ₄₀	Crystalline (bce)	Very low .	

water and protonated water are detectable, but for more dehydrated states they do not show any clear IR bands in contrast to silice or zeolite curalysts.

Recently, reliable IR spectra of FIFW-rHzQ with different edgrees of byteriation have been reported independently by upsilon and Zeochina's group. Fig. 7 shows our results where the water connects are quantified (reliable at least for n=0 and 6). Very broad bands ranging from 3500 to 1200 cm⁻¹ are evident. For n=6 the Assurption of the which the surveiture has been established, an extremely broad band is observed which we assigned to H_2 Q-(H_2 Q- H_1 - H_2 - H_1 - H_2 - H_2) greater in the hexaby-drate. A broad band having a peak at about 3200 cm⁻¹ was observed for 6 dehydrated sample (n=0–0.5), which was observed for 6 dehydrated sample (n=0–0.5), which was

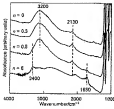


Fig. 7 IR spectra of $H_1PW_{12}O_{10}wH_2O$ with a variety of hydration states (0 < n < 6).

assigned to isolated acidic protons bonded to peripheral oxygens of the polyanion. These bands seem to exist in the IR spectra reported in earlier studies, but their presence and significance have not been well recognized.

Zecchina's group reported high quality IR spectra and provided valuable discussion of the IR banda.'1 The extremely broad band observed for the heaahydrate is reasonably interpreted by the nearly flat potential of a proton in a hydrogen bond. However, they possibly underestimated the water content of some samples (see below) and therefore their assignment of the IR bands is probably partially incorrect. The evacuation of hydrated HPVs at 300–340 K leads to the heaxhydrate which is fairly stable. See Usually a higher temperature (as high as co. 300 K) is necessary for further dehydration. Therefore, the sample of Zecchina's group which was obtained upon evocation at room temperature for 3 mins probably no 6, whereas the sample obtained by evacuation for 150 min is most likely the heaxhydrate.

The states and dynamics of protons and water in HPW have been clarified by an extensive study using solid-state 1¹H, 179 and 170 MAS NMR.¹²For example, the P NMR spectra of PWW-Hf₂O (n = 0.-6) which was prepared from the hexaly-drate by execution at 573-423 K are shown in Fig. 8. H NMR spectra measured at 288 and 173 K showed a very broad peak spectra measured at 288 and 173 K showed a very broad peak sharp peak for the other samples. The following important conclusions can be deduced from the NMR data.

(a) Acidic protons are present in three forms; (i) proton attached to polyanions, (ii) H₂O* (hydronium ion monomer) or H₂O strongly interacting with acidic protons, and (iii) H₂O*; (forthorium ion dimer), H₂O* and H₂O*, weakly interact with polyanions by hydrogen bonding. Acidic protons in the anhydrous sample attach to the polyanion leading to a salyalicant chemical shift in the ¹³P NMR spectrum. The probable bonding states are schematically illustrated in Fig. 9. On the basis of its studies, Zecchina's group concluded that

for intermediate hydration states (0 < n < 3), the acidic proton does not form k/O^2 but rather $O^2 + O^2 + O^2 + O^2$. Other, $O^2 + O^2 + O^2$. However, the close recomblance in the chemical shifts of the P NMR spectra (e.g. n = 2.1 measured at 173 K, Fig. 6) and of CoX⁴²²⁵ suggests the formation of H₂O² saddieg protons interact very weakly with polyanions. The remainder of the acidic protons remain directly bonded to the polyanions. It is should also be noted that the acid strength of HPW is greater than that of zeolites (see below, Table 3).

(b) The broad peak observed at 798 K (n = 0.5-4) splits imposeveral peaks at 173 K (Fig. 8, n = 0.5, 2.1 and 4.0). This means charither are of proton migration (exchange) is slow at 173 K and at 298 K is of the order of 200 Hz which is much faster than the rate of ordinary catalytic reactions. This study also indicated

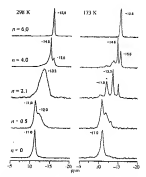


Fig. 8 ³¹P NMR spectra of H₂PW₁₂O₁₆nH₂O with a variety of hydration states (0 \leq n \leq 6) measured at 298 and (73 K.

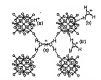


Fig. 9 Models proposed for the states of scielic protons and water in solid $H_1PW_{12}Q_{10}mH_2Q$ (0 < n < 6); two possible positions are shown for H_1Q .

Table 3 Acid strengths of various solid acids

Solid acid	Initial heat of NH, serption/kJ mol = 1	Approximate peak temporature of NH ₃ desorption/K
H ₂ PW ₍₂ O _{en}	195	850
Cso sH2 sPW12O40	170	830
SOJZIO1	165 (190)	1000 (as N ₂)
HZSM-5 (SVA) = 13)	150	670
SiO ₂ -Al ₂ O ₃	145	600
"Value in parentheses ref	brs to the very initial va	lue.

that water in the lattice enhances the mobility of protons. The high mobility of protons may facilitate protonation in acid catalysis.

(c) The relative intensities of the split peaks follow a binominal distribution, indicating a uniform (random) distribution of protons and water in the solid.

(d) In the anhydrous state (n = 0), there is one type of acidic proton species attached to the polyamion at least on the NMR time scale. This is consistent with our R data. We observed an apparently single broad band for the dehydrated sample (n =

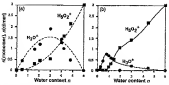


Fig. 10 Amounts of H₂O: (monomer) and H₂O₂: (dinter) per polyanism in $H_2PW_1/O_{ce}nH_2O(c) < n < 6$). (2) Samples prepared by evacuation of the heavily-tube at 173–420 K. Distinct lines were calculated by assigning that water molecules are randomly removed from the heavily-distinct. (b) Samples prepared to redwindsino of analysticus FFRW_1O₂M₂ in come targetimes.

(e) The concentrations of the three proton species in HPW-Mt-O (n = 0-6) change with the extent of dehydration as shown in Fig. 10(a). The trends are in agreement with those calculated by assuming random removal of water.

In contrast, HPW-nH₂O prepared by the addition of water to an anhydrous sample a room temperature is quite different [Fig. 10(b)]. The main reason for the difference may be the lower temperature of preparation for which the diffusion of water in slow. Rehydration leads mainly to H₂O₂ instead of H₂O₂ at a > 2. Careful preparations are always necessary to obtain HPA samples having uniform composition. In particular, special caution must be taken to zovid rehydration, since anhydrous HPAs are very sensitive to moisture (even from water adsorbed on wells of apparatus).

3.3 Acidity

The acid strength of HPAs vary in a wide range depending on the polyanion structure and its constitutuent elements (both hetero and addends atoms), as well as on the extent of hydration and reduction. Most results indicate that HPW after dehydration, the strongest HPA known so far, is a much stronger solid acid than zeolites including ZSM-5, and is close to that of superacids. 5.23 Indeed some believe HPW to be a superacid24 while others claim that its acidity is comparable with zeolites.21 The counter cation is also an important factor, Table 3 summarizes the acid strength as monitored by ammonia sorption (adsorption and absorption) and desorption for several solid acids,23,25 and Fig. 11 shows the results of calorimetric measurements of NH3 sorption on Cs2.5 and HPW at 423 K. Caution in interpretation of the results may be necessary for HPW which tends to exhibit pseudoliquid behaviour, Ammonia is absorbed into the bulk and forms ammonium salts, so that the lattice energy of the salts should be considered in calculating the heat of sorption.

Quantum chemical studies with higher levels of approximation on a full Keggin unit have recently been strempted. The stronger solidy of IFPW than H₂PNO₁₂O₂₀ (IFPMO) is indicated by density functional theory (DFT)-30 in early calculations, the position of protonation was interned to be at bridging oxygen calculation applied to a full Keggin unit including geometrical optimization.²⁵ Our ¹⁷O MAS NMR and IR data are in agreement with this conclusion.²⁵

There are a least five different mechanisms for the evolution of activity of solid IPAP satisfs* and hence the acticle properties are complicated unless the structures are carefully scrudnized. NMR satisfies revealed that protoco of a novel nature are formed by the reduction of Ag and Pd satis of HPM** Protocos are for the satisfies of the protocol of the protocol

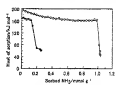


Fig. 13 Differential heats of NH₃ sorption measured at 423 K: (O) H₃PW₁₂O₄₀, (●) Cs_{2.5}H_{9.5}PW₁₂O₄₀.

the latter. The influence of the mobility of protons on catalytic performance is an interesting topic. Similar behaviour is observed for Ag zeolites and can be formulated by reactions (1)-(3).26°

$$2Ag^{+} + H_{2} \rightarrow 2Ag^{0} + 2H^{+}$$
 (1)

$$2Ag^0 + Ag^1 \rightarrow Ag_2^{-}$$
 (2)

$$Ag_5^+ + H_2 \rightarrow Ag_3H + H^+$$
 (3)

4 Recent topics in surface-catalysis of Cs and NH, salts of HPW

4.1 Microstructure

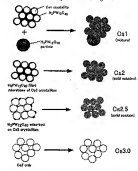
The microstructure, or the tentary structure (particle size, surface area, por distribution, e.c.) of insulvible scidic Ca and NH, salts of HPW is of great interest since Cd.5 showed extremely high catalytic activities; for various restolates in the liquid phase as a strong solid acid. ** "The activity is often more than ten times greater than that of zeolitics and more than there times that of the parent HPW. Activity much higher than that of TPPW was also reported for C2.4 be repeared in a slightly different manner. **D Furthermore, the size of the micropores of C2.4.5 in early uniform, can be controlled at the alm no level and leads to remarkable shape selective catalysis ^{2,33} Acidic NH, and are also active and for bemospherion (NH, HPPW ₁/Q₁) was more active than C_{2.2.5.73} Ag and TI salts secon to have stimilar microstructures.

The main reason for the high activity of Cs2.5 is its high surface acidity i.e. the large number of strongly acidic protons on the surface.3-* The number of surface sites is about half of the total number of protons contained in the solid owing to its high surface area (rea. 150 m² g "). Cs2.5 is a strong acid being only slightly weaker than the parent HPW (Table 3). Other

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reasons for the high activity are moderate hydrophobicity of the surface, a bimodal pore distribution and scid-base bifunctionality (see below). All of these factors are interesting and important for heterogeneous catalysis.

Extensive and detailed studies using ³¹P NMR, RRD, electron difficultion, AFM and ESM, adsorption of No, Ar, erc, as well as quantitative chemical analysis, have recently been carried out to clarify the mitorestrearce of the acidic Cs and NH, salts. ³⁵ Those measurements were applied compensively to the preparation processes i.e. solutions, precipitates and resulting solids obtained after drying and heat treatment, and provided a consistent view about the microtrustures of Cst.3 and other Cst salts. Their formation processes thus deduced are schemically illustractal in Fig. 12.



After drying After heat treatment
Fig. 12 Schematic illustration of the formation processes of acidic Cs salts:
Left; after drying, right; after heat treatment.

Upon tiration of an appecies solution of IPPW with a CssCO, agecues solution at 298 K, very fine precipitates of Cs3 (nanoparticles of cs. 10 mm in diameter) are formed, to which IPW is partly adopted with the remaining IPPW present in the solution. As the titration proceeds, the amount of IPPW in solution decreases, forming Cs3 precipitates. At the stoictionetry of Cs2 (CssCO, added: IPPW = 1:1), the precipitates are fine particles (allow cn. 10 mm in diameter) of Cs3. the surface of which are covered by nearly a monolayer of IPPW. Their surface area is very 10 wt 1 m² gr⁻¹), since the fine particles stoic together densely after drying. This model is supported by the fact that the particle size estimated from XDD investibits is co. 10 nm, while that estimated from the BET surface area is 500 –1000 nm.

When Cs₂CO₂ is added heyend a stolchlorotry of N = 2, the surface area increases sharply, since most of HPW pensipinates as Cd3 and the amount of HPW remaining in solution or adsorbed on precipitates diminishes rapidly. Hence, micropares start to develop which would have been absent if the remaining HPW had density connected the nanoparticles. ³²P SMR (which can differentiate between polyanious countaining 0, 1, 2 or 3 protons) demonstrated that thermit seatment at 373 473 K leads to a nearly uniform solid solution of Cs₂PW, ₂O₄₀ (CS3) and H₂PW, ₂O₄ (CHPW) will diffusion of protons and Cs' ions. the lattice constant changing accordingly. This process has been unambiguously confirmed by the formation of the identical solid solutions when Cs3, impregnated by various amounts of HPW, is treated at 473 K, as shown by the P NMR spectroscopy (Fig. 13).

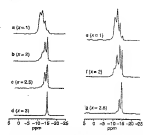


Fig. 13 Comparison of the ³¹P NMR spectra of Cb_2H3 , $_{sh}^{*}W_{12}O_{10}$ (CEX) [(a)+(d)] with Cs satis lawing the same compositions (CeX, X = 1, 2, 2.5) prepared by impregnation of $Cs_2PW_{12}O_{20}$, (C33) by HPW [(c)-(g)]. All samples were thermally (regted at 473 K.

C32.5 has a bimodal pore size distribution; micropores ranging from 0.5 to 1.0 nm (peak of 0.55 m and mmsty > 0.75 nm) and mesoperes (peak at 4.–5 nm) 3.64 lt is deduced that the micropores arise from spaces between nancorystallites (10–20 nm) in loose and random aggregates and mesopores arise from spaces between hancorystallites and between aggregates of size cn. 100–500 nm (see Fig. 2). The micropores account from about 70% of the total surface area of C4.2.5 (see below, Fig. 15(d)). Misfits in the connection of nanocrystallites have countly been suggested as the origin of small micropores (see below). Fig. 34 Another possible origin of micropores are polyanion vacancies as proposed for Cs.2 Micropore (no firm and detailed conclusion has been obtained for the origin of the micropores which have a runder sharp size distribution.

A surprising finding of the microstructure is the epitaxial assembly of Cs and NH₄ salts, which was discovered unexpectedly [Fig. 14 and Fig. 15].^{27,38} When the Cs salts are

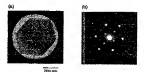


Fig. 14 Electron diffraction pattern of [NH₄):PW₁₂O₄₉ prepared by timution at 368 K using NH₄HCO₂: (a) the area in which the diffraction pattern was taken and (b) diffraction pattern.

prepared at a temperature as high as 370 K, the initially formed nanocrystallites (αa . 10 mm) assemble together with the identical orientation of crystal planes, leaving micropores between the nanoparticles [Fig. 15(b)], in contrast to Cs.2.5

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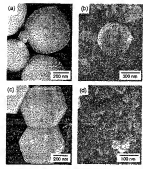


Fig. 15 SEM imagus of (a) (NH_a)₂PW₁₂O₄₀ prepared by titration at 273 K, (b) Cs₂PW₁₂O₄₀ (Cs₃) prepared by titration at 368 K, (c) (NH_a)₂PW₁₂O₄₀ prepared by titration at 368 K and (d) Cs₂PW₁₂O₄₀Cs₃) prepared by titration at 298 K, Cs₄S, is similar to Cs₃ in preparation.

prepared at room temperature [Fig. 15(d)]. This is most remarkable when a stoichiometric NH4 sali is prepared by means of homogeneous precipitation using urea decomposition.34 The initially formed particles are spherical and slightly oriented aggregates (resembling the NH₄ sait formed by titration at low temperature shown in Fig. 15(a)), that are comprised of nanocrystallites of size ca. 10 nm with micropores (ca. 0.6 nm) in the spaces between the nuncerystallites, and mesopores (2-10 nm) between the loosely bound nanocrystallites and/or between the aggregates (100-1000 nm). The particles then gradually turn into large 'crystalline' microporous aggregates (400-1000 nm) having regular dodecahedral shape and few mesopores [similar to but larger than the particles shown in Fig. 15(c)]. The particle size estimated by XRD linewidth measurements (>260 nm) is much greater than the size calculated from the surface area (10-20 nm), indicating that epitaxial connections occur between nanocrystallites. AFM and SEM images confirm that aggregates consist of fine particles of size ca. 10 nm. Electron diffraction (ED) shows regular discrete spots indicating crystallinity (Fig. 14), All of these observations indicate that the dodecahedral particles are 'crystalline' aggregates of nanoparticles and are porous. A monodispersed particle size (ea. 1000 nm) can be obtained by controlling the precipitation procedure (Fig. 16).30 Spherical aggregates shown in Fig. 15(a) and (b) also give discrete ED patterns.

4.2 Shape selectivity

Remarkable shape selectivity has been reported for serveral sacicalized reactions over C so also using molecules having catalyzed reactions over Cs so also using molecules having different sizes. While Cc2.5 is active for most reactions, Cc2.1 and Cc2.2 only catalyzed reactions of small molecules, 2^{6,42} Shape selective catalysis has also recently been found for conduction and bytrogenesation reactions. Occubers and cowerless or distance of the conduction of the comparison of the conduction o

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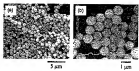


Fig. 16 SEM images of monedispersed (NH₄)₁PW₁₂O₈₀ prepared by iteration at 368 K

oxidation of methane, exrbon monoxide and benzage, and (ii) hydrogenation of relativine and cyclothecture. "Suppress executivity in the products was found for the reaction of n-hume," if yi has produced was found for the reaction of n-hume, "I The large-pore PC-S2.3 was very selective for the isomerization to isobatylene (94%), white cracked products (C_-C_) markedly increased with a decrease in the tisse of micropove (only 45% isobatene formed for Pr-(32.1) surshould to slow diffusion of the branched allocation in small micropoves (Octahara and connection of manocrystallities are the origin of fine micropoves of C2.2 (10.45 mm.)% as Illustrated in Fig. 17.

4.3 Surface acidity of Cs, H3-xPW12O49, CsX

The unusual change with. X of the statistic activity of CaX is reasonably interpreted by the unifies andily [Fig. 4(0)]. Mere, the surface activity is estimated by multiplying the number of polyunions and the surface (calculated from the surface area and the size of polyunion) and the number of protons per polyunion on the surface (from the proton content of the solid, or the chemical formula). Recently, the surface activity, or the number of protons, on the surface activity, or the number of protons, on the surface of CaX (X = 2.3), has directly been measured by an IR study of CO adsorption at I 10 X (Fig. 18). We can be considered to the control of CaX of the control of C

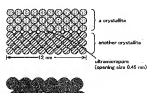
said a first CA strictural registion are adverted [Pig]. [3(0)].

As expected, the intensity of the first band (3165 cm⁻¹, CO).

As expected, the intensity of the first band (3165 cm⁻¹, CO) are as the said property of the strictural property of the strictural

4.4 Hydrophobicity: water-tolerant solid acid catalysts

It has been shown in an earlier study that the surface of organication of HPW exhibit hydrophobicity. P. Reutently, the hydrophobicity was semi-quantitatively evaluated for Cn2.5 and Cn3 by comparison of water and benzane adsorption. To bydrophobicity that evaluated is in the order of H2SM-5 (high silica) selfites or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Cx2.5 > Ex3SM-5 (two silica) > silica or Cn3 > Ex3SM-5 (two silica) > silica or Cn3 > Ex3SM-5 (two silica) > two silica > two silica



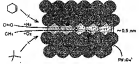


Fig. 17 A model of small micropores formed by misfits of two momory-stallines of Pt-Cs2, He-PW12Octo.

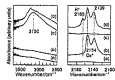


Fig. 18 IR spectrs of CO adsorbed on FFAs measured at 100 K. (a) Before and (b) after adsorption of CO on CspW12Oan (Cs3); (c) before and (d) after adsorption of CO on Cs2.xHa.yPW13Oan

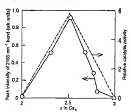


Fig. 19 Changes of the peak intensity of the 2165 cm⁻¹ band (CO advantor) on proton sites) and the estalytic activity of Cs,1]₁₋₁PW₂Co₄ (CsX) as a function of X for the rate of alkylation of 1.3.5-trimothylbename by cyclobexene at 353 K.

However, for the reaction of a nitrile and an alcohol in an excess of water, Cs2.5 was the most active as shown in Table 4.45

5 Efficient catalysts developed based on HPA

Table 5 lists recent examples of catalytic reactions using solid HPA. Earlier examples may be found in ref. 5(b).

Table 4 Comparison of catalytic activities of various solid solids for the reaction of corylnitrile and N-adamantanol to form N-adamantyleorylamide (NAA) in the presence of an excess of water.* Results in the absence of water shown in parentheses

Catalyst	Yield*(%)	Selectivity(%)	TON≠	
C8: sHasPW12O46	84 (97)	92 (93)	36 (42)	
HY zeolite	8 (79)	82 (89)	2 (0.2)	
Amberlyst 15	68 (100)	82 (81)	1 (1)	
Nation-H	77 (97)	84 (92)	6 (8)	
Nafinn-SiO+	40 (97)	94 (93)	22 (61)	

Rescribes conditions; catalyst: 0.2 g. ceryltitrile: 60 mand, 1-edestrationol: 1.3 mmd, 375 K, 6 h. 5% Yield; 100 y. NAA formed/(Yedemantanol) added), 4% Selectivity: 100 y. QNAA formed/(Yedemantanol) added), 4% Selectivity: 100 y. QNAA formed/(YAA) formed resyltation formed). "TON (tupnover number); mol NAA formed/mol aeld sites in catalysts.

5.1 Bifunctional catalysts

HPAs show acidity as well as unique basicity, these properties as well as their oxidizing ability can be controlled over a wide range which is of use in cashyst design. The co-existence of these properties can be utilized to prepare bifunctional and multifunctional catalysts. It has been shown that the oxidation of methacrolical to methorcylic acid proceeds in two steps; the first step is soid-oasilyzed exterification to from an intermediate and the second step the oxidation of the intermediate which is rest-efermining. Hence this reaction can be catalyzed in a interesting to not that these two years of the properties of the controlled interesting to not that these two proceedings of the properties and of oxidation of isobutyric acid to enablespile acid, with acidity accolerating side-fractions (Scheme 2).

The much higher activity of Cs2.5 than conventional solid actifs can not be explained by actific properties alone [Fig. 3(c)], so that actid-base bifunctional catalysis was suggested for Cs2.5.5%

Efficient catalytic reactions can be realized by the combination of FIPA canalysas with noble meals. One-stage oxidation of ethylene to acedic acid has been commercialized (10000 ton yr-") by combining a Keggain-ype IPA cetalyst and Pd.⁴⁶ Here, the addition of Se or Te to Pd is casential to supports the Here, the addition of Se or Te to Pd is casential to supports the suggested to proceed in two steps (e.gu. (4) and (3)). a Westertype mechanism (via acetaldehyda) being excluded. Control of existing a support of the processing the step of the catalyst

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Table 5 Acid-catalyzed reactions with solid beteropoly compounds

Reaction	Cmalyst	Reference
$RCO_2H + R'OH \rightarrow RCO_2R'$	H ₃ PW ₁₂ O ₄₆ ,	59
Alkylation of alkylphenol by isobutylene (shape selective)	H ₂ SiW ₁₂ O ₄₆ /MCM-41 H ₂ PW ₁₂ O ₄₆ /MCM-41	53 <i>b</i>
. Trioxene + phenol	Silica-included	60
1 Transle - Micitar	H ₂ PW ₁ C ₂₀	00
Isobutane + n-butanes Cs alkylates	H-PW - O - MCM-41	61
	K2 H42PW12O4n (supercritical)	62
Diels-Alder reaction of oulnoac	H.PW. Oas	63
Acylation of gylene	Cs-HPW; Oct	530
Adamantylamide synthesis	Cs2 sHa sPW tr Oan	45
Hydration of directly/butene	Cs2 dla PW (2Oan	44a
n-C, iso-C,	(NHL,Cs.H),PW13O4	64
	Pt-Cs2,4H0 +PW12O40	
Oxidation of othylene to acetic acid via ethanol	Pd(To)-SiW12O40	46
	(commercialized)	
Ethyl acotate from socile acid	HPA (commercialized)	
Michael addition	H.PW, zOir. (pseudoliquid)	160
$E:C(CH_2OH)_2CH_2OCH_2O(CH_2OH)_3Et + H_2O \rightarrow$	H.PW12O40, etc.	i4
ZExC(CH2OH)2CH2OH + HCHO	In new phase?)	
These reactions are mostly taken from a list produced by Professor T. Okuban		

CH1
CH-mc-CRO
Southern-A-cididy
Congressive
ACH-COQH
Stilt-yr-(II)
Olidative r.A-cididy
Congressive
CH-co-COQH
Stilt-yr-(II)
Olidative r.A-cididy
CRO-CH-COQH
Stilt-yr-(III)
CRO-COQH
Stilt-yr-(III)
Scheme 2
Socheme 2
Socheme 2

development. (The photo on the cover is the plant used for this process developed by Showa Danko, Co., Ltd.)

C-H-+ H-O -> EtOH (acid catalysis) (4)

 $E(DH + O_1 - MeCO_2H + H_2O)$ (conclusions) (5) $E(H + O_1 - MeCO_2H + H_2O)$ (conclusions) (5) $E(H + O_1 - MeCO_2H + H_2O)$ (conclusions) (6) the combination of Pd or Pt with Ca2.5 creates active and selective catalyses for the isomerization of r-allanes $E(C_1 - C_2)^{-1/2}$. Descrivation and creaking which are significantly observed over Cs2.3 alone are dramatically diminished by the addition of mobile metals in the presence of H_2 . Without addity, the activity

is very low. Interestingly, acidity completely prevents the hydrogenolysis of alkanes extratived by noble metals. A mechanism essentially based on a classical bifunctional entalysis has been prosposed, and canalyzed insemiration of ethydrogenation-bydrogenation. Recently, a mechanistic study using 20 Labeled n-buranes* has revealed that the selective situmerization to isobutane over Pt dispersed on Cs.2.5 mostly takes place by a monomolecular mechanism, while the reaction proceeds less selectively u/a s bimolecular mechanism in the case of Cs.2.5 alone (Scheme 3). At higher reaction temperatures, the

contribution of the bimolecular mechanism increases also for Pt-Cs2.5. Kozhevnikov and coworkers reported that HPW

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combined with Pd produces only soft coke in the oligomorization of propene and the removal of coke by combustion takes place at a much lower temperature than in the case of HPW alone which forms both soft and bard cokes.⁴⁹

5.2 Supported HPA catalysts

Hydrogen forms (or free acids) of HPAs assually have low surface areas. On the other hand, very active Ca2. Suring a surface areas on the other hand, very active Ca2. Suring a large nirface area tends to become a mility inspension during liquid-phase reasotions, which makes it difficult to separate the cazalyst after the reaction is finished. To solve these problems, many attempts have been made to disperse and fits. HPA catalysts on various supports, where the ashelity of the HPA and firm fixation are the key issues. Support materials such as silice, carbon, and organic mains have been applied with varying levels of success, with new supporting materials such methods being actively pursued. Although the seventure and composition of supported HPAs are somatines uncertainly incatalytic activities are often observed and separation made

Izumi et al, prepared HPW and Cs2.5 included in a silica matrix by an in situ sol-gol method. Recently, shape selectivity was observed for alkylation of phenol by formaldehyde owing to the micropores of a silica matrix. 50 Supported HPAs prepared by this method have been applied as photocatalysts.51 Soled et al. reported an in situ preparation of Cs2.5 inside silica particles.52 Cs-containing silica is added to an aqueous solution of HPA. As the HPA solution diffuses into porce of silica, a Cs salt starts to precipitate at a certain level of concentration. resulting in an egg-yolk type imprognation. Large-pore zeolites like MCM-4153 and layered clays have been applied as supports of HPA. HPAs loaded in layered double hydroxide were active for epoxidation and showed shape selectivity 54 In situ synthesis of HPAs in the supercage of Y-zeolite is an interesting method for the preparation of supported HPA catalysts.35 For metal oxide supports, interactions between the surface and the HPA often degrade the polyanion structure. On basic solids such as MgO and Al2O3, the Keggin structure readily decomposes, as expected from instability of HPAs in aqueous solution at high pH. Even on the surface of silica, which has only weak interactions with HPAs, these tend to decompose to smaller clusters. In most cases the decomposition is significant at a low loading level whereas the starting polyanion structure is predominant when the loading level is high. Supporting on or imbedding in organic polymers has also been attempted. Recent examples include polyazamethines, 26st polyaniline,366 and polyphenylene oxide.36r As expected from the moderate stabilities of organic ammonium and oxonium salts, the HPA structures appear to remain mostly intact. In addition, chemical interactions between the polymer and HPA sometimes modify the catalytic performance in a desirable manner

6 Future

Notable procress has been achieved recently in heterogeneous acid catalytic reactions of HPAs. Full utilization of the pseudoliquid phase and further development of bifimetional and shapescientive catalysis will be interesting targets for the future. Specifically organized secondary and tertiary structures that are synthesized by using novel cations and polyanions may open up new areas of catalysis. If the structure, composition and stability of polyanions are properly controlled on supports, solid HPA catalysts will find much wider practical applications. The development of regeneration methods for deactivated HPA cetalysts is another important subject for practical applications. As for catalysis in solution, unconventional reaction fields such as multi-phase catalysis are promising. From the viewpoint of fundamental study of HPA catalysis, the basicity of the surface of the heteropolyanion (or unique complexation character) together with its role in catalysis is of interest and understanding of the catalytic reaction at the molecular/atomic level is expected to be accomplished in the near future.

More progress is anticipated for oxidation catalysis of HPAs although oxidation catalysis lies outside the scope of the present article. The design of primary structures (structure and composition) has been successful for exidation in solution and may be extended to heterogeneous catalysis, if HPAs are stabilized or reaction systems chosen carefully. Examples include diiron and dimanganese substatuted Keggin anions as shown in Fig. 1(c)57 which efficiently catalyze selective oxidation of alkanes, although enhancement of reaction rate is still desirable. A variety of polyanion structures and compositions (new and known) as well as recent progress in novel synthetic methods promises the development of efficient catalysts based on HPAs. For example, an exotic HPA synthesized by Newmann and Dahan is efficient in selective oxidation, 5x Owing to various advantages, HPAs are hoped to play important roles as green catalysts in chemical syntheses in a sustainable manner.

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